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The role of global dietary transitions for safeguarding biodiversity

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1 **The role of global dietary transitions for safeguarding biodiversity.**

2
3

4 **Abstract**

5

6 Diets lower in meat could reduce agricultural expansion and intensification thereby reducing
7 biodiversity impacts. However, land use requirements, associated with alternate diets, in
8 biodiverse regions across different taxa are not fully understood. We use a spatially explicit
9 global food and land system model to address this gap. We quantify land-use change in locations
10 important for biodiversity across taxa and find diets low in animal products reduce agricultural
11 expansion and intensity in regions with high biodiversity. Reducing ruminant meat
12 consumption alone however was not sufficient to reduce fertiliser and irrigation application in
13 biodiverse locations. The results differed according to taxa, emphasising that land-use change
14 effects on biodiversity will be taxon specific. The links shown between global meat consumption
15 and agricultural expansion and intensification in the biodiverse regions of the world indicates
16 the potential to help safeguard biodiverse natural ecosystems through dietary change.

17 **Key words:** biodiversity, land use change, diet, consumption

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19

20

1. Introduction

Expansion of agricultural land, together with intensified management practices are some of the greatest threats to the conservation of terrestrial ecosystems and biodiversity (Machovina et al., 2015; Machovina and Feeley, 2014; Marchal et al., 2011; Newbold et al., 2015; Ripple et al., 2014a). Over 35% of the Earth's permanent ice-free land surface is currently used for food production (Foley et al., 2005), with the expansion of agricultural land for food production in the last 300 years having reduced natural grasslands by up to fifty percent and natural forests by one third (Goldewijk, 2001). The associated loss of natural ecosystems has had negative consequences for biodiversity (Gibson et al., 2011; Pereira et al., 2012; Pimm et al., 2014). Agricultural intensification that increases yields can reduce the area of land needed for production, but can also harm biodiversity through fertiliser and pesticide pollution (Flohre et al., 2011; Gibbs et al., 2009; Kleijn et al., 2009) as well as irrigation abstraction on ecological river flows (De Frutos et al., 2015; Yamaguchi and Blumwald, 2005). Land-use change models have demonstrated that biodiverse regions will be significantly threatened by future agricultural expansion and intensification (Delzeit et al., 2017; Kehoe et al., 2017, 2015). Protected areas can be an effective contribution to prevent agricultural expansion (Pringle, 2017), but conservation efforts that focus on food demand will also play a role.

Meat production has been associated with higher land and water use, and higher GHG emissions, per unit of energy or protein than other foods (Machovina et al., 2015; Poore and Nemecek, 2018; Tilman and Clark, 2014a). In particular, heavily managed and densely stocked pastures pose serious threats to biodiversity (Machovina and Feeley, 2014; Ripple et al., 2014a). 65% of agricultural expansion in recent decades has been associated with increased production of animal products (Alexander et al., 2015), and land-use changes associated with animal husbandry account for roughly 30% of current global biodiversity loss (Westhoek et al., 2011). Livestock production is increasing most rapidly in tropical regions with high biodiversity (Machovina et al., 2015). The tropics are also experiencing the highest rates of species extinction (Dirzo et al., 2014), at a time when global extinction rates have been estimated to be 1000 times the geological background rate (Pimm et al., 2014, 1995). Much future human population growth is expected to occur in these biodiverse tropical nations, and as incomes continue to rise in developing countries, animal product consumption is expected to increase further (Machovina et al., 2015; Stoll-Kleemann and Schmidt, 2017). If current trends in animal product consumption continue, and if industrialised countries do not reduce high rates of meat consumption, it is estimated that one billion additional hectares of natural land will be cleared for agriculture by 2050 (Tilman et al., 2011, 2001).

Reducing meat consumption would not only improve global human health—consumption of meat in industrialised countries is currently double the amount that is deemed healthy (Wellesley et al., 2015)—but the Intergovernmental Panel on Climate Change (IPCC) also identified it as an important focus for climate change mitigation (Pachauri et al., 2014). Modelling studies have quantified land-use changes associated with dietary shifts, demonstrating that demand-side reductions in meat consumption could reduce GHG emissions and deforestation (Bajželj et al., 2014; Erb et al., 2016; Popp et al., 2010; Stehfest et al., 2009; Tilman and Clark, 2014a; Wirsenius et al., 2010). However, fewer studies (Kok et al., 2018; Tilman et al., 2017; Visconti et al., 2016) have considered the effects of diet on biodiversity, and

none have explored the spatial impacts across multiple taxa. The spatial nature of biodiversity and variations in distributions between taxa means that spatially explicit analyses are required to understand the impact of dietary choices on biodiversity.

Here we address this critical gap in understanding the environmental consequences of food-system changes. We use a global food-system model (PLUMv2/LPJ-GUESS, Alexander et al., 2018) to explore land use and agricultural intensity change until 2100 under three alternative dietary scenarios: Business-as-usual (BAU), 95% reduction in ruminant product consumption (LOW-R), 95% reduction in animal product consumption (LOW-AP). This work is unique in considering the spatially disaggregated consequences of future dietary scenarios for high biodiversity locations across different taxa. We also, for the first time, consider the role of nitrogen and irrigation intensity changes on locations important for biodiversity.

2. Methods

2.1. Modelling framework

PLUMv2 is a global land use and food-system model that combines spatially-explicit, biophysically-derived yield responses with socio-economic scenario data to project future demand, land use, and management inputs (Alexander et al., 2018). For each country and time-step, the agricultural land use and level of imports or exports is determined through a least-cost optimisation that meets the demand for food and bioenergy commodities in each country. Food demand is projected based on log-linear relationships with per-capita income using GDP and populations from the Shared Socioeconomic Pathway (SSP) scenarios (O'Neill et al., 2014). Demand for food and bioenergy commodities is projected at a country level for six commodity groups: cereals, oilcrops, pulses, starchy roots, ruminant products, and monogastric products. Demand for dedicated energy crops (i.e., second-generation bioenergy) is specified as a global trajectory with all production locations determined endogenously. Food and bioenergy demand is met by in-country expansion or intensification of crops or from imports from the global market. ~~Over~~ Production of commodities in ~~excess of a~~ country's domestic demand are exported to the global market. ~~In PLUMv2 supply and demand in t~~ The global market is not constrained to ~~be in~~ equilibrium, ~~with~~ over- or under- supply of commodities ~~can be~~ buffered through explicitly modelled stocks. ~~For each commodity a single tariff free price exists in each time step and the initial price was set exogenously but subsequently adjusted in each of the following time periods according to under- or oversupply in the global market. Prices are updated for the next year based on the aggregate imbalance of imports and exports in that year. For example over supply of a commodity on the global market decreases the price; this reduces the benefits from its export and reduces the cost of importing it, creating a tendency to correct for the oversupply. For each commodity a single tariff free price exists in each time step, which is adjusted for transport costs and other barriers, e.g. tariffs, to obtain country specific prices. For example over supply of a commodity decreases the price; this reduces the benefits from its export and reduces the cost of importing it, creating a tendency to correct for the oversupply.~~

Crop yield responses used in PLUMv2 are provided on a 0.5° grid by a dynamic global vegetation model, LPJ-GUESS (Smith et al., 2014), for a range of fertilisation rates and rain-fed vs irrigated conditions. Other management practices (e.g. pesticide application, machinery stock, reseedling of grassland) are represented in PLUMv2 by a "management intensity" factor. Natural land cover here is comprised of forested primary land, non-forested primary land and abandoned agricultural land. ~~In the grid cells four decision variables (i.e. area, fertiliser,~~

Commented [HR1]: In general I've added to the methods section to address the following comment

'The authors refer to a recent model development paper but to stand at its own more detailed information is needed.'

Commented [HR2]: REVIEWER COMMENT: How is trade dealt with within the modelling framework? It is mentioned in line 93-4 but not expanded on.

115 irrigation and other intensity) for each of the eight land use types (seven crop types plus
116 pasture) are determined during the optimisation, resulting in more than 77,000 decision
117 variables in each year. To achieve this PLUMv2 uses the spatially specific crop yield responses
118 to intensity inputs, various land use costs, such as land conversion costs and input costs, and
119 trade costs. This ultimately determine land use solutions that meet country level demand.

Commented [AP3]: What 'this'? Just "PLUMv2 uses..."?

Commented [AP4]: Again, unclear what 'this' refers to.

Commented [HR5]: Reviewer comment: **Line 309-311**
"plum chooses not to abandon existing cropland" – is
this a results or is PLUMv2 hard-coded into the model
to do this?

121 Socioeconomic parameters are in line with the "middle of the road" SSP scenario (SSP2), which
122 also provided population and GDP trajectories (Dellink et al., 2017; Jones and O'Neill, 2016). The
123 SSPs describe alternative global societal pathways through the 21st century (O'Neill et al., 2015,
124 2014). SSP2 is the middle of the road pathways with trends largely exhibiting historic patterns.

125 Population and GDP trajectories were taken from SSP2 using World Bank projections (IIASA,
126 2014), and demand for food commodities were taken from FAOSTAT (FAOSTAT, 2015a, 2015b).
127 The climate and atmospheric CO₂ forcing scenario RCP 6.0 was used as it is considered the
128 Representative Concentration Pathway (van Vuuren et al., 2011) most consistent with SSP2
129 (Engström et al., 2016). Forcings were taken from the 1850–2100 IPSL-CM5A-MR outputs from
130 the Fifth Coupled Model Intercomparison Project (CMIP5; 71). First- and second-generation
131 bioenergy demand trajectories are specified exogenously to represent a business-as-usual
132 scenario with no specific climate change mitigation policies. Demand for first-generation
133 bioenergy is modelled from an observed baseline level of demand (Alexander et al., 2015;
134 FAOSTAT, 2015a) adjusted to double by 2030 from the 2010 level and thereafter remain
135 constant. Global demand for dedicated second-generation bioenergy crops increases to 4000 Mt
136 DM/year by 2100, in line with the SSP2 demand with baseline assumptions (Popp et al., 2016).
137 For parameter settings that were not specified exogenously from available data on existing and
138 future trends, for example technology change rates, expert judgement was used to align
139 quantitative parameter settings with the qualitative SSP2 storyline. Scenario elements of the
140 SSP2 narrative that were assumed to influence changes in the PLUMv2 input parameters were
141 identified. Qualitative changes in parameters were estimated based on an interpretation of the
142 SSP2 storyline (Engström et al., 2016). These qualitative estimates of parameters and
143 uncertainty levels were translated into quantitative values characterised by a uniform
144 distribution. Each parameter therefore had a range defined by a range of 50% above and below
145 the central parameter values. A Monte Carlo approach to explore uncertainty associated with
146 input parameters was used and parameters were sampled using a Sobol sequence method with
147 n = 30 (Chalaby et al., 2015); the central parameter values used in each of the scenarios can be
148 found in Appendix C, Table C2. This approach allowed us to capture the uncertainty within the
149 model framework.

150

151 2.2. Protected areas

152 The proportion of protected land with a status of "designated" and IUCN category I–VI within a
153 grid cell is calculated using the WDPA database (UNEP-WCMC., 2016). Within each grid cell,
154 natural land designated as protected is prevented from conversion to any form of agricultural
155 use. Within each grid cell, a minimum fraction (5%) of primary unprotected natural land is also
156 prevented from agricultural use due to assumed limits to agricultural production, e.g. field
157 boundaries, roads/tracks, and other farm infrastructure. Slope constraints (IIASA/FAO, 2010)
158 also prevent agricultural use in regions of high altitude. In cases where agricultural land already
159 exceeds the area specified as protected, no further agricultural expansion can occur. China's
160 National Forest Protection Program is implemented as an annual limit to deforestation of 1.1%
161 in China. (Ren et al., 2015).

162

163

2.3. Scenario description

2.3.1. Business as usual (BAU)

This scenario assumes that the shift in consumption away from staples, such as pulses and starchy roots, and towards animal products continues as incomes rise. The relationship between rising income and increasing consumption of commodities such as meat, milk, and refined sugars has been observed historically and therefore in line with the SSP2 pathway, we assume future consumption trends in the baseline largely exhibit historic patterns (Keyzer et al., 2005; Tilman et al., 2011).

2.3.2. 95% reduction in ruminant product consumption (LOW-R)

This scenario represents a major shift in world consumption patterns of ruminant products, this could be potentially driven by increasing meat prices induced by stricter climate and health policies, consumer awareness and increasing land and animal feed expenses. This scenario assumes that the consumption of ruminant products decreases steadily from 2010 to 2100 until crop products replace 95% of ruminant product consumption. Ruminant products are replaced by a mixture of cereals, starchy roots, pulses, and oilcrops; however, the same calorie intake is maintained. An example of the dietary changes in terms of per capita consumption and the proportions of substitution and are given in Appendix C, Table C1 and Table C2 respectively. 95% was chosen for the stylised scenarios to demonstrate the potential effects a very large, but not total, reduction in ruminant product consumption would have on global land use.

2.3.3. 95% reduction in animal product consumption (LOW-AP)

This scenario is similar to the above, but assumes that non-meat commodities replace both ruminant and monogastric consumption. As with the LOW-R scenario 95% was chosen for to demonstrate the potential effects a very large, but not total, reduction in animal product consumption would have on global land use.

2.4. Exploring the consequences of dietary change for biodiversity

2.4.1. Conservation International (CI) biodiversity hotspots

The 35 CI hotspots cover 2.3% of the land surface but support 50% of the world's endemic plant species and 43% of vertebrate endemic species. To qualify as a hotspot, a region must be threatened —i.e. contain at most 30% of its original natural vegetation—yet contain at least 1500 endemic vascular plants. The CI biodiversity hotspot database is used to identify particular regions of importance for biodiversity (Mittermeier et al., 2004; Myers et al., 2000). CI hotspot shapefile data are converted to 0.5° raster maps. Any 0.5° cell containing CI hotspot polygon data is classified as a CI hotspot irrespective of hotspot size. The CI map is therefore binary and cells are classified as either a CI hotspot or not.

2.4.2. Vertebrate species richness maps

Criteria for the biodiversity hotspots database only account for vascular plant species richness. Thus, we also consider maps of vertebrate species richness, small-range vertebrate species richness, and threatened species richness (Jenkins et al., 2013; Pimm et al., 2014). The resolution of the vertebrate species richness maps was decreased from 0.1° to 0.5° resolution to match PLUMv2; the mean species richness was calculated for each grid cell. For all taxa, the distribution of species richness across grid cells is right-skewed: most cells contain a few species while there are a few cells with a large number of species. For each taxon's map we therefore convert the mean species richness values of grid cells into percentile values

Commented [AP6]: Might read better for each of the scenarios to phrase first sentence as.

"The business-as-usual scenario ..." or equivalent.

Not essential

Commented [HR7]: REVIEWER COMMENT: On the scenario description (section 2.3) it would be good to have more contextual information used in the justification of the settings chosen for the modelling. Were they taken from existing trends or entirely taken from the expert judgment of the modelling team. To what extent are the settings chosen plausible?

I have also added a section to the results/discussion about limitations related to this.

(richness index). We assume that ‘species-rich regions’ comprise cells with a richness index ≥ 0.9 , i.e. the 90th percentile of grid cells and therefore, similar to the CI hotspots, we focus on those regions with the greatest biodiversity (appendix A, figures A1-3).

We explore land use change, agricultural expansion, and intensification projected by PLUMv2 in CI hotspots and in vertebrate species-rich regions for the different dietary scenarios. We consider the loss of natural land, forests, and natural grasslands and changes in input intensities such as fertiliser and irrigation in grid cells classed either as CI hotspots or with a richness index ≥ 0.9 . From this, we identify regions of threat using a threat index: regions with high biodiversity that overlap with areas of projected agricultural expansion. We calculate this overall threat index for all species in each 0.5° grid cell. This is the proportion of natural land projected to be lost by 2100 multiplied by the summed richness index of birds, mammals, and amphibians for the median PLUMv2 parameter simulation run. For the threat index we therefore assume each species is equally important regardless of taxon.

3. Results and discussion

3.1. Land cover change in biodiverse regions

In agreement with results from previous modelling studies (Delzeit et al., 2017; Kehoe et al., 2017, 2015), the most threatened regions—locations with high biodiversity under pressure from agricultural expansion—are in the tropics under BAU scenarios (Figure 1). Scenarios of lower animal product consumption (LOW-R and LOW-AP) greatly reduce agricultural expansion in regions of high biodiversity compared to the BAU scenario (Figure 1 & 2). By 2100, 9% (984 Mha) of global natural land is lost, of which 95% is in the tropics—equivalent to 24% of natural land in these latitudes (Figure 2). Conversely, reduced animal product consumption (LOW-AP) resulted in a 7% (703 Mha) increase in global natural land between 2010–2100 (Figure 2, Appendix B Figure B1) with lower losses across the tropics (Figure 1, Figure 2) and increases in natural land across the northern hemisphere (Figure 2). Deforestation and land clearing for agriculture have been identified as the leading causes of biodiversity decline (Gibson et al., 2011). Therefore, the potential for dietary change to reduce global agricultural expansion by approximately 1687 Mha (11% of global land area) is an important finding for biodiversity conservation (Laurance et al., 2012; Pereira et al., 2012).

Species-rich regions across the different taxa are largely found in the tropics (Appendix A, Figure A1-3) and the greatest loss of natural land in species-rich regions occurs in BAU (Figure 3). In BAU, on average, 98% of global pasture expansion takes place in the tropics as demand for ruminant products in tropical countries increases with increasing population and income (Appendix C, Figure C2). As incomes increase, consumption shifts from staples such as starchy roots and pulses to commodities such as meat, milk, and refined sugars (Keyzer et al., 2005; Tilman et al., 2011). However, the rate of increasing consumption of animal products slows and plateaus with any further rise in income (Cole and Mccoskey, 2013), which is also represented in the log-linear relationships with per-capita income used in our model (Alexander et al., 2018). Consequently, in developing tropical countries, the transition from low incomes to high incomes results in greater demand of ruminant products (Appendix C, Figure C2), and pasture expands at the expense of natural land. In contrast, income and the animal product consumption in developed countries outside the tropics are already high, with large areas of existing pasture meeting demand for ruminant products. Given the relationship between income and consumption, increases in income in developed countries do not lead to further large increases in demand for animal products (Appendix C, Figure C2). Under LOW-AP and LOW-R,

260 abandonment of existing pasture in developed countries leads to large increases of natural land
261 at a global level (Figure 2). This does not coincide with large increases in natural land in
262 species-rich regions (Figure 3), however, because it largely takes place in locations that are not
263 here classified as species-rich — i.e. those outside the tropics. This result can be seen when
264 comparing Figures A1–A3 in Appendix A (spatial distribution of species-rich regions for the
265 different taxa) with Figure 2. In species-rich regions the LOW-R and LOW-AP scenarios reduce
266 pasture expansion rather than increase natural land. Although existing pasture in the tropics is
267 also abandoned, this is offset by cropland expansion (see below); therefore natural land area in
268 species-rich regions under LOW-R and LOW-AP is relatively stable, compared to BAU (Figure 3).

269 Cropland expands by 28% in the tropics under BAU to produce crops for food and as animal
270 feed as demand for animal products grows in the developing world. Under LOW-R, cropland
271 expands by 38% in the tropics, this is greater than under BAU because demand for food and
272 feed for monogastrics is the same while additional crops are required to replace ruminant
273 products. Under LOW-AP, despite reduced demand for feed for animals, existing cropland area
274 in 2010 is not sufficient to produce enough crops to replace animal products and meet food
275 demand of a growing population; consequently, cropland still expands by 27%. The greater
276 cropland requirements under LOW-R explain the marginally greater losses of natural land in
277 species-rich regions (Figure 3) in LOW-R compared to LOW-AP. However, on average, the
278 amounts of water and nitrogen applied to cropland in the tropics under LOW-AP are 42% and
279 68% less, respectively, than under BAU. Therefore, while the total area of cropland remains the
280 same in the tropics, the intensity of agricultural inputs declines under LOW-AP with the
281 reduction in demand for animal feed.
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283
284

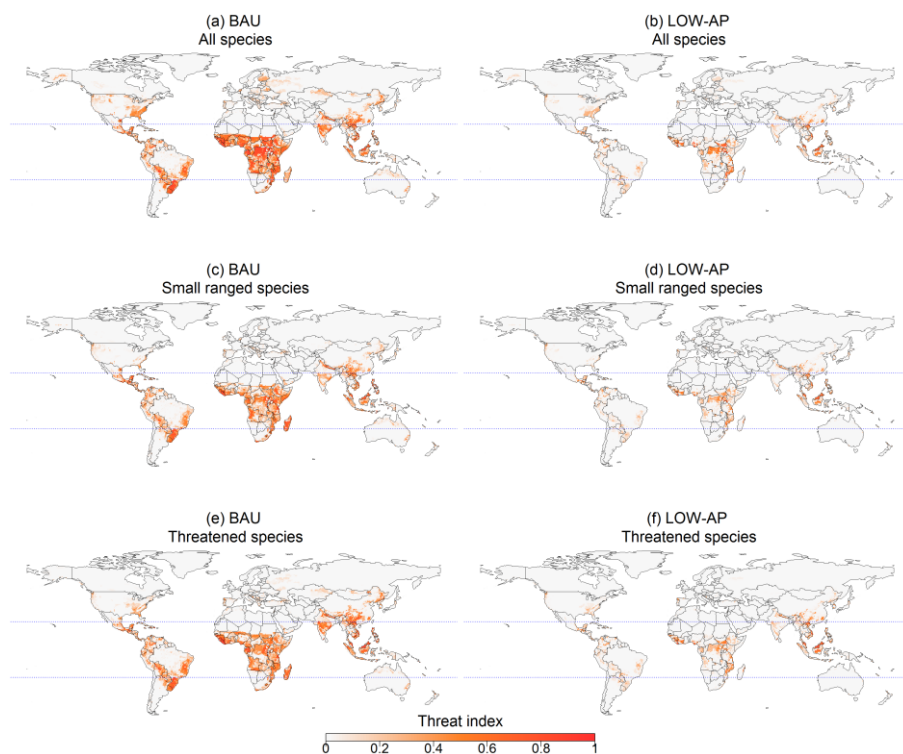


Figure 1: Spatial distribution of regions of threat; regions with high biodiversity under pressure from agricultural expansion. The left column (a,c,e) is the BAU scenario and the right column (b,d,f) is the LOW-AP scenario for the different types of species richness. Blue dashed lines delineate the tropics.

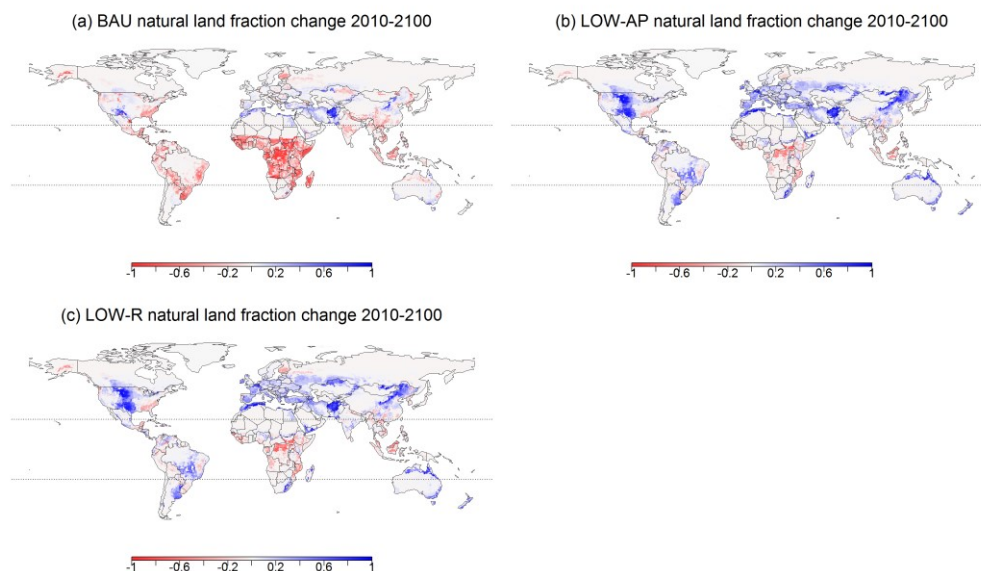


Figure 2: Change in natural land cover fraction between 2010-2100 for (a) BAU (b) LOW-AP (c) LOW-R. Dotted lines delineate the tropics.

Tilman et al., (2017) investigated the biodiversity of mammals and birds in a non-spatial, country-level approach and found dietary change reduced extinction risk. However, previous spatially explicit studies typically only consider single taxon with amphibians particularly underrepresented. For example, Visconti et al., (2016), considering only mammals, found consumption change could reduce extinction risk. The locations classified here as species-rich differ between mammals, birds and amphibians (see Appendix A, Figures A1–3). Differences regarding the impacts of land-use change therefore arise between and within taxa, and are important when considering conservation targets (Ceballos and Ehrlich, 2006; Jenkins et al., 2013; Orme et al., 2005; Pimm et al., 2014; Possingham and Wilson, 2005). For example, natural land loss in parts of Ecuador overlap to a greater extent with regions of threatened bird species-richness than with regions of threatened mammal-species-richness. While broad patterns are similar across CI hotspots and taxa—for example, the greatest loss of natural land occurs in BAU while the LOW-R and LOW-AP scenarios result in smaller losses or increasing natural land cover (Figure 3)—important differences remain. For example, with LOW-AP, while the small ranged species-rich regions shows little change or decreases in natural land cover by 2100, the grid cells within the threatened species-rich region show increasing natural land cover by 2100. Changing dietary patterns may therefore have the greatest benefits for threatened species in term of habitat recovery. Measures of total species richness are important when considering threats to overall range size and ecosystem functioning related to population sizes. However, the richness of small-ranged species and/or threatened species are often regarded as more appropriate measures when planning conservation to prevent extinctions (Ceballos and Ehrlich, 2006). Visconti et al., (2015), for example, highlighted the importance of considering the status of taxa from a protected area perspective: Targeting protection towards threatened species had positive effects on suitable habitat for terrestrial mammals, while expanding protected areas according to ecoregion targets had negative effects. Furthermore, a number of studies have demonstrated that hotspots of species richness between taxa and classifications of taxa are often incongruent (Jenkins et al., 2013; Orme et

Commented [HR8]: REVIEWER COMMENT: The directions of the trends discussed aren't really surprising, but perhaps there is something in the magnitudes or spatial patterns that could be picked out to highlight the novelty more keenly?

Commented [AP9R8]: Hard to know what's been added and therefore to assess how well it deals with the point raised.

al., 2005; Pimm et al., 2014). The differences between CI hotspots and type of taxa here further support the argument that no single metric is sufficient when considering threats to, and the conservation of, biodiversity (Ceballos and Ehrlich, 2006; Jenkins et al., 2013; Orme et al., 2005; Pimm et al., 2014; Possingham and Wilson, 2005).

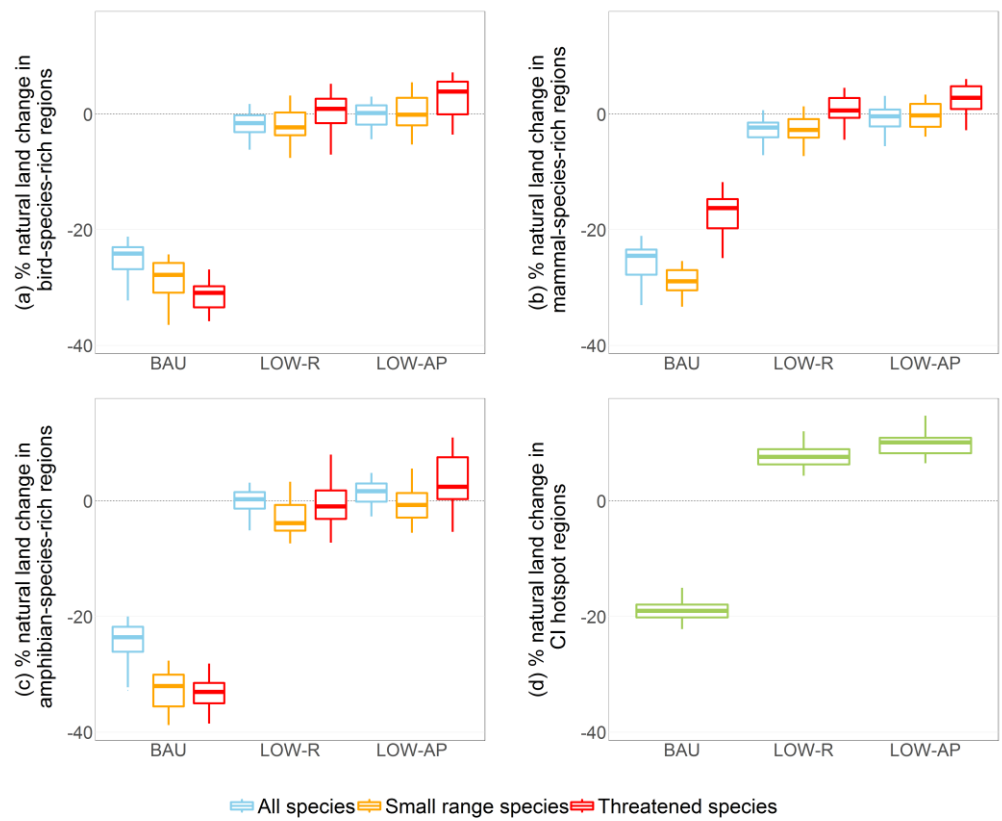


Figure 3: Projected natural land change by 2100 in (a) bird-, (b) mammal-, and (c) amphibian-species-rich regions, and (d) CI hotspots for the different dietary scenarios. Species-rich regions are comprised of cells with a richness index ≥ 0.9 . Colours in a–c represent the different types of species-rich regions: all species (blue), small-ranged species (orange), and threatened species (red). Boxplots distributions generated with $n=30$.

334 3.2. Agricultural intensity change

335 The type and level of agricultural management has an important role in the impact on different
 336 taxa, for example, at a European scale Flohre et al., (2011) found that while agricultural
 337 intensity negatively affected the species richness of birds it did not affect carabid beetles (De
 338 Frutos et al., 2015; Flohre et al., 2011; Gibbs et al., 2009; Kleijn et al., 2009; Yamaguchi and
 339 Blumwald, 2005). *It has been shown that* increasing nitrogen, in particular, reduces plant
 340 biodiversity (Bobbink et al., 2010; Reich, 2009; Stevens et al., 2004) with consequences for
 341 faunal biodiversity (Nijssen et al., 2017). *In their review* Bobbink et al., (2010) highlighted that
 342 the negative effects of nitrogen accumulation on biodiversity has occurred across a wide range
 343 of ecosystems and geographic areas. Nijssen et al., (2017) *recently* identified ten pathways
 344 through which increased nitrogen alters faunal biodiversity. N-driven faunal decline has been
 345 demonstrated, for example, in some rare bird species where elevated nitrogen reduced
 346 vegetation heterogeneity and/or preferred habitat with consequences for prey abundance (de
 347 Vries et al., 2011). However, *while pathways identified by Nijssen et al., (2017) were supported*
 348 *by peer reviewed literature* there remains knowledge gaps regarding the mechanisms that drive
 349 observed biodiversity changes. *In terms of* Similarly, irrigation there is a *wide* body of evidence
 350 that demonstrates the negative effects of water extraction on natural ecosystems, *with and*
 351 *agricultural water use is one of the leading causes of the majority of* freshwater withdrawal *in*
 352 *the world globally used in agriculture*. The disruption of water flows and river regulation has, for
 353 example, altered floodplain forests resulting in their dieback globally. Such forests are
 354 ecologically important due to their high biodiversity, *with and* climate change induced droughts
 355 are likely to further exacerbate forest mortality (Horner et al., 2009). In Spain, the expansion of
 356 irrigated agriculture has coincided with the disappearance of up to 61% of biodiverse wetlands
 357 over the last fifty years (Fuentes-Rodríguez et al., 2013). Furthermore, intensive livestock
 358 farming that involves irrigation has also been found to substantially alter water chemistry of
 359 nearby rivers with potential consequences for both aquatic and riparian species diversity
 360 (Martin-Queller et al., 2010). Given the *widespread* implications *for biodiversity* of increasing
 361 nitrogen and irrigation use *for biodiversity* the need to consider such consequences are
 362 apparent. However, no previous land use modelling studies have explored changes in irrigation
 363 and nitrogen fertiliser intensities that are associated with reductions in meat consumption in
 364 biodiverse regions. Agricultural intensity is typically represented in land-use models in a
 365 stylised and spatially aggregated manner, making the evaluation of their impacts challenging
 366 (Lotze-Campen et al., 2008; Nelson et al., 2014). Our analysis addresses this gap and by
 367 including spatially specific crop responses to different inputs in our modelling framework, *W*
 368 *we* are able to show the effects of dietary changes on *intensity* inputs *intensity, with a focus on in*
 369 *species-rich regions where biodiversity impacts are likely to be most acute.*

370 In the LOW-R scenario, demand for monogastric feed crops is unchanged from BAU, while
 371 demand for food crops increases to replace ruminant products (Appendix C, Figure C1). This net
 372 increase in crop demand results in crop area expansion relative to BAU. However average global
 373 fertiliser and irrigation intensities (on a per-area basis) change similarly to BAU; the median
 374 increase in nitrogen and irrigation, in CI hotspots and species-rich regions, under LOW-R are
 375 similar to BAU (Figure 4, Figure 5). LOW-AP decreases demand relative to BAU for
 376 monogastrics, as well as ruminants, and consequently decreases demand for crops as feed
 377 (Figure C1). Rather than reduce cropland area this results in reduced nitrogen and water inputs
 378 in these locations. From a fertiliser and irrigation perspective, reduced feed production
 379 therefore has the greatest potential to reduce inputs and replacing pasture-fed ruminant
 380 products alone may not have substantial benefits for biodiversity.

381

Commented [AP10]: Compared to what else?

Commented [AP11]: Two sentences of the form "A said <something>". Could at least one, perhaps both be rephrased so avoid that rather boring structure.

Commented [AP12]: Delete? Seems to add little.

Commented [AP13]: 70% from memory!? Worth a reference too. Aquastat?

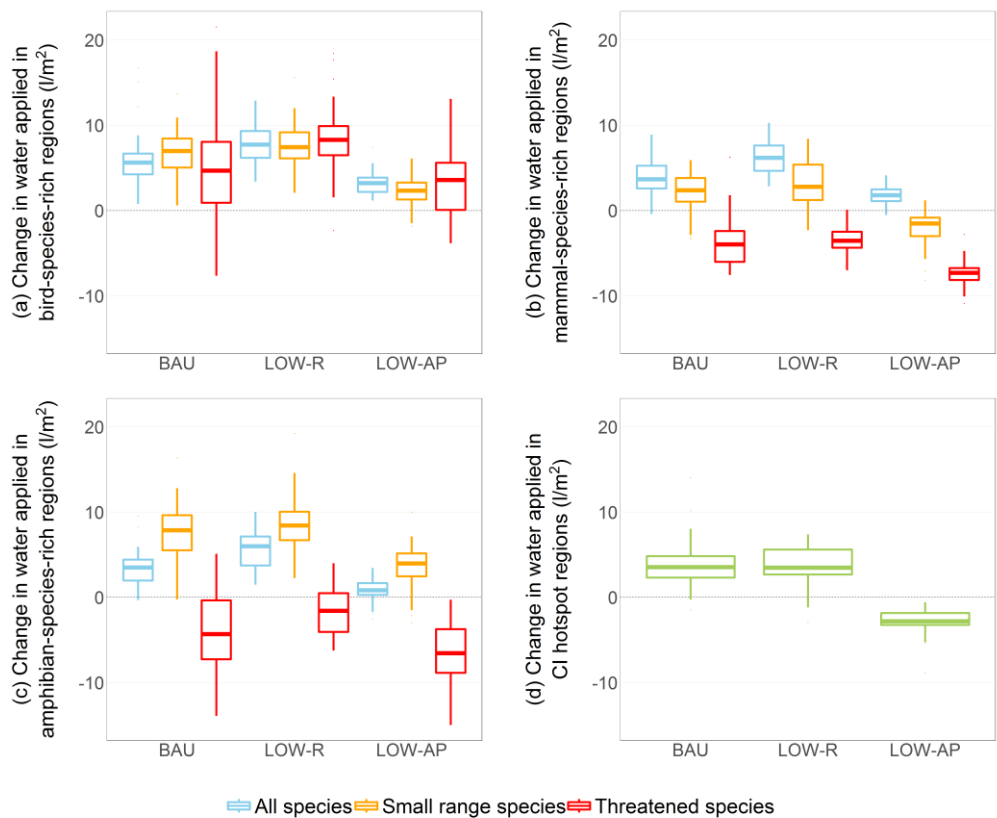
Commented [AP14]: Right word? "retreat" "reduction in area". Might need a reordering of phrase.

Commented [AP15]: Rather weak evidence! Sure lots of things have coincided with that. Coincided with increased call for Catalanian independence as well!

Commented [HR16]: REVIEWER COMMENT: The link between nitrogen application and biodiversity is clear, but could also do with a little more justification. The link between irrigation and biodiversity is less clear-cut. What are the thoughts / concerns here?

382 The differences in nitrogen and water inputs between alternate dietary scenarios highlight the
 383 need to consider fertiliser and irrigation individually. Any potential intensity changes associated
 384 with dietary change will require scrutiny as measures to reduce agricultural expansion may not
 385 necessarily reduce intensification. We do not consider a scenario with a 'livestock revolution'—
 386 a shift away from pasture-based production toward industrialised production that requires
 387 crop-based feeds (Delgado et al., 2001; Naylor et al., 2005; Swain et al., 2018)—which could
 388 similarly reduce the rate of agricultural expansion, but with increased intensity. There is an
 389 inherent trade-off between agricultural intensification and expansion. Intensification is more
 390 polluting, but requires less land, while expansion is less polluting, but requires more land.
 391 Ultimately, both can have negative consequences for biodiversity and thus managing this trade-
 392 off is complex. For example, the recent IPBES Regional Assessment for Europe & Central Asia
 393 recommends that Europe reduce agricultural intensity to conserve European biodiversity
 394 (IPBES, 2018). However, this could displace food production and the associated consequences
 395 for biodiversity, through imports, to other parts of the world.

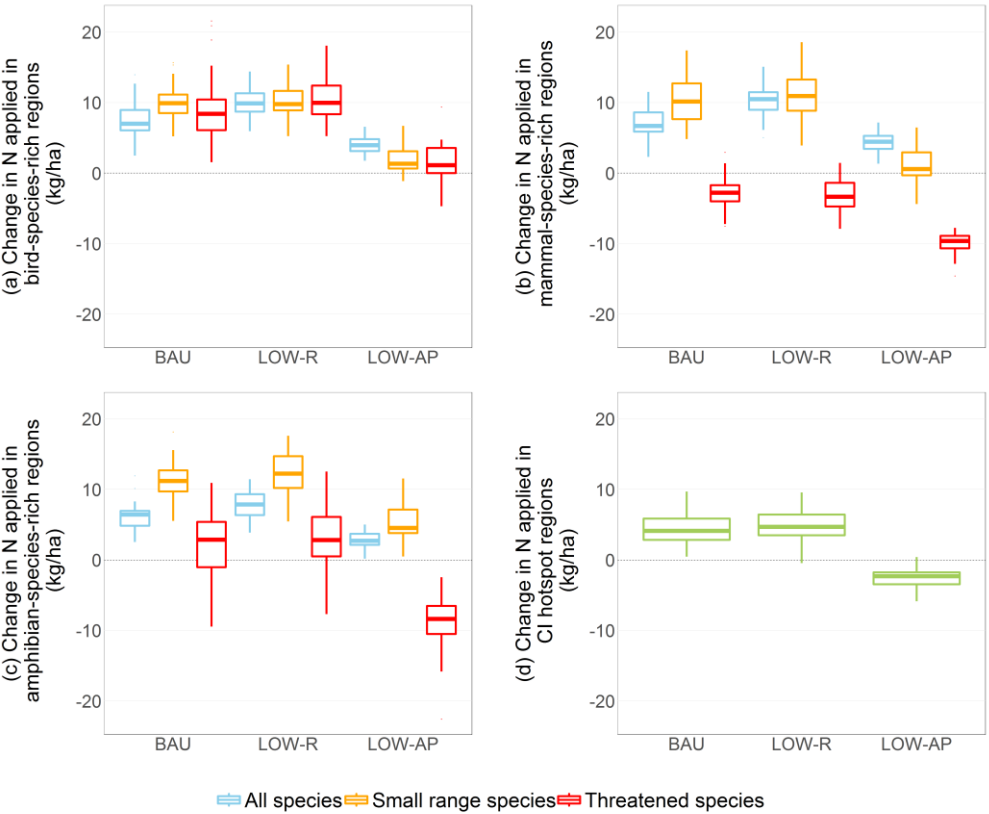
396



397

398 *Figure 4: Projected change in irrigation water use by 2100 in (a) bird-, (b) mammal-, and (c)*
 399 *amphibian-species-rich regions and (d) CI hotspots for the different dietary scenarios. Species-rich*
 400 *regions are comprised of cells with a richness index ≥ 0.9 . Colours in a–c represent the different*

401 types of species-rich regions: all species (blue), small-ranged species (orange), threatened species
 402 (red). Boxplots distributions generated with n=30.
 403



404
 405 *Figure 5: Projected nitrogen fertiliser intensity change by 2100 in (a) bird-, (b) mammal-, and (c)*
 406 *amphibian-species-rich regions and (d) CI hotspots for the different dietary scenarios. Species-rich*
 407 *regions are comprised of cells with a richness index ≥ 0.9 . Colours in a–c represent the different*
 408 *types of species-rich regions: all species (blue), small-ranged species (orange), threatened species*
 409 *(red). Boxplots distributions generated with n=30.*

410 Within scenarios the intensity results show large differences across species-rich regions,
 411 establishing the need to consider land expansion jointly with land management when assessing
 412 biodiversity impacts of land-use change, and to provide these analyses for individual taxa of
 413 different status. For example, increases in irrigation water applied in locations rich in small-
 414 ranged amphibians are greater compared to locations rich in small-ranged birds or mammals
 415 (Figure 4). Without separating out taxa, such a finding could be overlooked, despite the
 416 probable greater importance of irrigation water withdrawal for amphibian populations. The
 417 intensity change results are heterogeneous between the different regions of species richness
 418 because food demand, the crops grown and yield response to agricultural inputs are location-

specific (see methods). We find, for example, nitrogen and irrigation application in bird-species-rich regions increases over the period 2010–2100 (Figure 5a). Conversely, nitrogen and irrigation application declines in threatened mammal- and amphibian-species-rich regions. In the threatened mammal and amphibian locations by 2100 under BAU agricultural area expands (Figure 3) and consequently agricultural production has increased sufficiently to meet demand such that less nitrogen and water are required. In the LOW-AP scenarios, in the threatened mammal and amphibian locations, reduced nitrogen and water use reduces with shrinking agricultural area (Figure 3). Changing dietary demand may therefore have the greatest benefits for threatened species through the reduction of both agricultural land area and agricultural inputs in regions of high biodiversity.

Commented [AP17]: Not sure I understand. Or the next sentence.

Commented [HR18]: REVIEWER COMMENT: One striking finding is that threatened species behave markedly different in Figure 4 and 5 – it would be worth investigating this more. Why is this the case and what are possible conservation implications here?

3.3. Uncertainty and limitations

The stylized scenarios here assume high substitution rates of animal products, 95%, similar to other studies that have assumed shifts towards complete vegetarianism (Stehfest et al., 2009; Tilman and Clark, 2014a) or large reductions, e.g. Visconti et al., (2016) assumptions imply reducing meat and egg consumption in all regions by 76–88%. Such scenarios are useful for illustrating the effects of dietary transitions on land use changes, however, arguably such large scale shifts will face barriers as dietary choices that individuals make are influenced by a number of factors such as culture, price, availability, taste and convenience. Taking such factors into account may reduce the potential for large scale dietary change. Lower rates of animal product substitution would inevitable result in lower environmental benefits in this study and others. For example, Stehfest et al., (2009) found a healthy diet that included some level of animal product consumption resulted in greater land use and GHG emissions than scenarios that reduced meat or animal product consumption entirely.

Commented [HR19]: REVIEWER COMMENT: Also, a more critical discussion of underlying uncertainties would be very useful. How robust are these results and how would the biodiversity implications change if the spatial patterns looked different?

REVIEWER COMMENT: There is more space for a discussion of the modelling as a whole within the broader context of the literature and particularly for some reflection on the limitations of the approach (especially the impacts of input choices on final results).

REVIEWER COMMENT: I suggest its publication, after minor revision. In particular I don't find clearly an estimation of the uncertainties linked to the models used for the analysis.

Commented [AP20]: Long and hard to follow sentence

We explore land use change in regions with the greatest levels of biodiversity by include CI hotspots and grid cells that are in the 90th percentile for species richness in our analysis. Our regions of interest are therefore largely have a focused in the tropics. Using absolute species richness loss has the advantage of highlighting particularly biodiverse regions at risk, with land use change in these areas will therefore potentially have having a disproportionate effect on global biodiversity loss. Similar to other studies we find these highly diverse regions, such as sub-Saharan Africa and Latin America, are suitable for large scale agricultural expansion further highlighting their importance in terms of conservation. However the choice to focus on areas with the highest biodiversity inevitably means land use change and the consequences for temperate or other climate zones are not well addressed in this study here. Changing dietary patterns will undoubtedly affect such regions too for example reducing animal product consumption reduced natural land loss in the south east of the United States, Kazakhstan and Mongolia (Figure 2). However, while these regions do not necessary harbour high levels of biodiversity, and are therefore not in our 90th percentile, they nevertheless may contain species of significant conservation or cultural importance. Therefore while it was out with the scope of our study to consider all geographic regions the consideration of temperate zones should not be overlooked.

Commented [HR21]: REVIEWER COMMENT: On the scenario description (section 2.3) it would be good to have more contextual information used in the justification of the settings chosen for the modelling. Were they taken from existing trends or entirely taken from the expert judgment of the modelling team. To what extent are the settings chosen plausible?

Commented [AP22]: Reading this paragraph makes is seem like only the tropics is included. That's not right either. Ok, so the tropics is over-represented, but don't think it's quite as extreme as implied overall in this text (perhaps I'm wrong through).

Commented [AP23]: Would it not be good to explicitly make the point here that we are considering changes in diets globally (including from these temperate regions) even those the focus is on the biodiversity in tropical regions.

Commented [HR24]: REVIEWER COMMENT: The choice of focusing on only areas in the top 10% of species richness numbers is not quite clear. This way, large (e.g. virtually the whole temperate zone) areas are excluded ex ante from further analysis. Obviously changing diets will also have a huge impact on them. It would be nice to see this also addressed in the study.

Land use changes to meet demand in PLUMv2 arise through a complex decision making process that involves assessing spatially explicit crop yield responses, a variety of agricultural costs and trade related costs. The parameter settings used in this study produced benchmarking results in line with historical data (Alexander et al., 2018). However agricultural costs may change with

future economic development and policies. While we use a Monte-Carlo approach to incorporate uncertainty of the input parameters, rates of agricultural expansion would reduce if we increased the cost of agricultural expansion. Similarly if agricultural input costs, such as fertiliser, were to rise the intensity of fertiliser use would decrease. Both outcomes would likely have beneficial effects for biodiversity in the BAU scenario, thereby lessening the magnitude of area and intensity savings with dietary change. The assumptions regarding future socio-economic and climate condition as based on SSP2 and RCP 6.0 respectively. Further analysis under a range of SSP trajectories may alter the land use patterns we find. For example changing GDP, population size or political shifts, such as increasing protectionism under SSP3, would alter the baseline food demand projections and change food supply requirements with consequences for land use. SSPs that therefore project more demand than SSP2 used here, particularly in developing tropical countries, will likely result in greater agricultural expansion in biodiverse regions. Similarly alternative climate pathways may have consequences for projected intensity use in biodiverse regions. Increased atmospheric CO₂ levels are linked to higher yield potentials, reduced nitrogen losses and greater water use efficiency. In previous work this leads to lower fertiliser and irrigation inputs in PLUMv2 (Alexander et al., 2018). Therefore, while lower climate forcing's could be beneficial for climate change, they may have unexpected negative effects such that more intensity inputs are required in agriculture to achieve desired yields to meet demand. Global agricultural cropland projections could also potentially shift as cropland area in regions negatively affected by climate change is reduced while cropland area in regions with increased crop potential grows. Changing both the SSP and RCP trajectory used in modelling studies of biodiversity may therefore alter the spatial patterns of threats to biodiversity. Indeed recent modelling studies comparing SSPs and RCPs found that stronger mitigation scenarios, corresponding to lower RCPs, had greater benefits for biodiversity (Chaudhary and Mooers, 2018; Newbold et al., 2015).

3.4. Conclusions and perspectives

We find diets low in animal products reduce agricultural expansion and intensity in regions with high biodiversity and the magnitude of change differed according to taxa, emphasising that land-use change effects on biodiversity will be taxon specific. Numerous tropical countries with high biodiversity have rates of increasing per capita meat production, and several are projected to require up to 30% more agricultural land by 2050 (e.g. Ecuador, Brazil, and China) (Machovina et al., 2015). Our results also demonstrate the importance of developing countries, particularly those in the tropics, for biodiversity. The transition from low incomes to high incomes and the associated increase in animal product consumption in developing countries drives large losses of agricultural land across the tropics and in species-rich-regions under BAU. In many developing countries, access to sufficient protein is limited and demand-side measures such as global dietary interventions could be detrimental to the welfare of populations and thus not ethical. Efforts to preserve biodiversity and ecosystem functioning will therefore require scrutiny to ensure that changes are complementary to food security goals in developing countries (including nutritional requirements) and respectful of cultural heritage. Land use change in our BAU scenario is comparable to socio-economic conditions within the shared socio-economic pathway (SSP) scenario SSP2 ('middle of the road', Popp et al., 2016). Likewise, the scenarios of reduced meat consumption have been uniformly applied across countries. Future scenarios of land-use change associated with alternative diets should encapsulate aspects of fairness and equity (Tilman and Clark, 2014b). For example, a reduction of animal product consumption in developed countries combined with the sustainable trade of meat into countries with animal-protein deficits could simultaneously increase the health of industrialised countries and prevent the destruction of natural land in tropical regions.

Dietary change, will be most effective if implemented as part of a suite of demand-side and supply-side measures to reduce biodiversity loss (Tilman et al., 2017; Visconti et al., 2016). In a modelling approach that combined increasing vegetarianism with reduction of food waste, by 2030, agricultural land decreased to a greater extent than we find here under LOW-AP (Wirsén et al., 2010). Policy screening scenarios similarly found that reaching any biodiversity target will require a combination of strategies: for example, dietary change combined with waste reduction and more efficient agricultural practices (Marchal et al., 2011; Ten Brink et al., 2010). Reducing global meat consumption, and other demand-side measures such as reducing food waste, will be socially and politically complex. It has been suggested that large-scale dietary change will require incentives or regulations (Ripple et al., 2014b). Furthermore, global diet alterations will need to complement food security goals and address global food inequalities. However, biodiversity is an essential component of ecosystem functioning, as well as human well-being, e.g. via provisioning of ecosystem services (IPBES, 2018; Naeem et al., 2016). Efforts to preserve biodiversity are, therefore, of the utmost importance and may require dietary change.

4. References

- Alexander, P., Rabin, S., Anthoni, P., Henry, R., Pugh, T., Rounsevell, M., Arneth, A., 2018. Adaptation of global land use and management intensity to changes in climate and atmospheric carbon dioxide. *Glob. Chang. Biol.*
- Alexander, P., Rounsevell, M.D.A., Dislich, C., Dodson, J.R., Engström, K., Moran, D., 2015. Drivers for global agricultural land use change: The nexus of diet, population, yield and bioenergy. *Glob. Environ. Chang.* 35, 138–147. doi:<http://dx.doi.org/10.1016/j.gloenvcha.2015.08.011>
- Bajželj, B., Richards, K.S., Allwood, J.M., Smith, P., Dennis, J.S., Curmi, E., Gilligan, C.A., 2014. Importance of food-demand management for climate mitigation. *Nat. Clim. Chang.* 4, 924–929. doi:10.1038/nclimate2353
- Bobbink, R., Hicks, K., Galloway, J., Spranger, T., Alkemade, R., Ashmore, M., Bustamante, M., Cinnerby, S., Davidson, E., Dentener, F., Emmett, B., Erisman, J.-W., Fenn, M., Gilliam, F., Nordin, A., Pardo, L., De Vries, W., 2010. Global assessment of nitrogen deposition effects on terrestrial plant diversity: a synthesis. *Ecol. Appl.* 20, 30–59. doi:10.1890/08-1140.1
- Ceballos, G., Ehrlich, P.R., 2006. Global mammal distributions, biodiversity hotspots, and conservation. *Proc. Natl. Acad. Sci.* 103, 19374–19379. doi:10.1073/pnas.0609334103
- Chalaby, Y., Dutang, C., Savicky, P., Wuertz, D., 2015. Toolbox for Pseudo and Quasi Random Number Generation and RNG Tests Version. Comprehensive R Archive Network (CRAN).
- Chaudhary, A., Mooers, A., 2018. Terrestrial Vertebrate Biodiversity Loss under Future Global Land Use Change Scenarios. *Sustainability* 10, 2764.
- Cole, J.R., Mccoskey, S., 2013. Does global meat consumption follow an environmental Kuznets curve? *Sustain. Sci. Pract. Policy* 9, 26–36.
- De Frutos, A., Olea, P.P., Mateo-Tomás, P., 2015. Responses of medium- and large-sized bird diversity to irrigation in dry cereal agroecosystems across spatial scales. *Agric. Ecosyst. Environ.* 207, 141–152. doi:<https://doi.org/10.1016/j.agee.2015.04.009>
- de Vries, W., Erisman, J.W., Spranger, T., Stevens, C.J., van den Berg, L., 2011. Nitrogen as a threat to European terrestrial biodiversity, in: Sutton, M.A., Howard, C.M., Erisman, J.W., Billen, G., Bleeker, A., Grennfelt, P., Van Grinsven, H., Grizzetti, B. (Eds.), *The European Nitrogen Assessment: Sources, Effects and Policy Perspectives*. Cambridge University Press, pp.

565 436–494.

566 Delgado, C., Rosegrant, M., Steinfeld, H., Ehui, S., Courbois, C., 2001. Livestock to 2020: the next
567 food revolution. *Outlook Agric.* 30, 27–29.

568 Dellink, R., Chateau, J., Lanzi, E., Magné, B., 2017. Long-term economic growth projections in the
569 Shared Socioeconomic Pathways. *Glob. Environ. Chang.* 42, 200–214.

570 Delzeit, R., Zabel, F., Meyer, C., Václavík, T., 2017. Addressing future trade-offs between
571 biodiversity and cropland expansion to improve food security. *Reg. Environ. Chang.* 17,
572 1443. doi:10.1007/s10113-016-0944-0

573 Dirzo, R., Young, H.S., Galetti, M., Ceballos, G., Isaac, N.J.B., Collen, B., 2014. Defaunation in the
574 Anthropocene. *Science* (80-.). 345, 401 LP-406.

575 Engström, K., Olin, S., Rounsevell, M.D.A., Brogaard, S., Van Vuuren, D.P., Alexander, P., Murray-
576 Rust, D., Arneth, A., 2016. Assessing uncertainties in global cropland futures using a
577 conditional probabilistic modelling framework. *Earth Syst. Dyn.* 7, 893–915.
578 doi:10.5194/esd-7-893-2016

579 Erb, K.-H., Lauk, C., Kastner, T., Mayer, A., Theurl, M.C., Haberl, H., 2016. Exploring the
580 biophysical option space for feeding the world without deforestation. *Nat. Commun.* 7.

581 FAOSTAT, 2015a. Commodity Balances/Crops Primary Equivalent (2015-12-16). Food and
582 Agriculture Organization of the United Nations, Rome, Italy.

583 FAOSTAT, 2015b. Commodity Balances/Livestock and Fish Primary Equivalent (2015-12-16).
584 Food and Agriculture Organization of the United Nations, Rome, Italy.

585 Flohre, A., Fischer, C., Aavik, T., Bengtsson, J., Berendse, F., Bommarco, R., Ceryngier, P., Clement,
586 L.W., Dennis, C., Eggers, S., Emmerson, M., Geiger, F., Guerrero, I., Hawro, V., Inchausti, P.,
587 Liira, J., Morales, M.B., Oñate, J.J., Pärt, T., Weisser, W.W., Winqvist, C., Thies, C., Tschardtke,
588 T., 2011. Agricultural intensification and biodiversity partitioning in European landscapes
589 comparing plants, carabids, and birds. *Ecol. Appl.* 21, 1772–1781. doi:10.1890/10-0645.1

590 Foley, J.A., DeFries, R., Asner, G.P., Barford, C., Bonan, G., Carpenter, S.R., Chapin, F.S., Coe, M.T.,
591 Daily, G.C., Gibbs, H.K., 2005. Global consequences of land use. *Science* (80-.). 309, 570–
592 574.

593 Fuentes-Rodríguez, F., Juan, M., Gallego, I., Lusi, M., Fenoy, E., León, D., Peñalver, P., Toja, J., Casas,
594 J.J., 2013. Diversity in Mediterranean farm ponds: trade-offs and synergies between
595 irrigation modernisation and biodiversity conservation. *Freshw. Biol.* 58, 63–78.
596 doi:10.1111/fwb.12038

597 Gibbs, K.E., Mackey, R.L., Currie, D.J., 2009. Human land use, agriculture, pesticides and losses of
598 imperiled species. *Divers. Distrib.* 15, 242–253.

599 Gibson, L., Lee, T.M., Koh, L.P., Brook, B.W., Gardner, T.A., Barlow, J., Peres, C.A., Bradshaw, C.J.A.,
600 Laurance, W.F., Lovejoy, T.E., 2011. Primary forests are irreplaceable for sustaining
601 tropical biodiversity. *Nature* 478, 378.

602 Goldewijk, K.K., 2001. Estimating global land use change over the past 300 years: The HYDE
603 Database. *Global Biogeochem. Cycles* 15, 417–433. doi:10.1029/1999GB001232

604 Horner, G.J., Baker, P.J., Mac Nally, R., Cunningham, S.C., Thomson, J.R., Hamilton, F., 2009.
605 Mortality of developing floodplain forests subjected to a drying climate and water
606 extraction. *Glob. Chang. Biol.* 15, 2176–2186. doi:10.1111/j.1365-2486.2009.01915.x

607 IIASA, 2014. SSP Database (version 0.93). International Institute for Applied Systems Analysis,
608 Laxenburg, Austria, International Institute for Applied Systems Analysis, Laxenburg,

609 Austria.

610 IIASA/FAO, 2010. Global Agroecological Zones (GAEZ v3.0).

611 IPBES, 2018. Summary for policymakers of the regional assessment report on biodiversity and
612 ecosystem services for Europe and Central Asia of the Intergovernmental Science-Policy
613 Platform on Biodiversity and Ecosystem Services. IPBES secretariat, Bonn, Germany.

614 Jenkins, C.N., Pimm, S.L., Joppa, L.N., 2013. Global patterns of terrestrial vertebrate diversity and
615 conservation. *Proc. Natl. Acad. Sci.* 110, E2602–E2610. doi:10.1073/pnas.1302251110

616 Jones, B., O'Neill, B.C., 2016. Spatially explicit global population scenarios consistent with the
617 Shared Socioeconomic Pathways. *Environ. Res. Lett.* 11, 84003.

618 Kehoe, L., Kuemmerle, T., Meyer, C., Levers, C., Václavík, T., Kreft, H., 2015. Global patterns of
619 agricultural land-use intensity and vertebrate diversity. *Divers. Distrib.* 21, 1308–1318.
620 doi:10.1111/ddi.12359

621 Kehoe, L., Romero-Muñoz, A., Polaina, E., Estes, L., Kreft, H., Kuemmerle, T., 2017. Biodiversity at
622 risk under future cropland expansion and intensification. *Nat. Ecol. Evol.* 1, 1129–1135.
623 doi:10.1038/s41559-017-0234-3

624 Keyzer, M. a., Merbis, M.D., Pavel, I.F.P.W., van Wesenbeeck, C.F. a., 2005. Diet shifts towards
625 meat and the effects on cereal use: can we feed the animals in 2030? *Ecol. Econ.* 55, 187–
626 202. doi:10.1016/j.ecolecon.2004.12.002

627 Kleijn, D., Kohler, F., Baldi, A., Batary, P., Concepcion, E., Clough, Y., Diaz, M., Gabriel, D.,
628 Holzschuh, A., Knop, E., Kovacs, A., Marshall, E.J., Tschardtke, T., Verhulst, J., 2009. On the
629 relationship between farmland biodiversity and land-use intensity in Europe. *Proc. R. Soc.*
630 *B Biol. Sci.* 276, 903–909. doi:10.1098/rspb.2008.1509

631 Kok, M.T.J., Alkemade, R., Bakkenes, M., van Eerd, M., Janse, J., Mandryk, M., Kram, T., Lazarova,
632 T., Meijer, J., van Oorschot, M., Westhoek, H., van der Zagt, R., van der Berg, M., van der Esch,
633 S., Prins, A.-G., van Vuuren, D.P., 2018. Pathways for agriculture and forestry to contribute
634 to terrestrial biodiversity conservation: A global scenario-study. *Biol. Conserv.* 221, 137–
635 150. doi:https://doi.org/10.1016/j.biocon.2018.03.003

636 Laurance, W.F., Useche, D.C., Rendeiro, J., Kalka, M., Bradshaw, C.J.A., Sloan, S.P., Laurance, S.G.,
637 Campbell, M., Abernethy, K., Alvarez, P., 2012. Averting biodiversity collapse in tropical
638 forest protected areas. *Nature* 489, 290.

639 Lotze-Campen, H., Müller, C., Bondeau, A., Rost, S., Popp, A., Lucht, W., 2008. Global food demand,
640 productivity growth, and the scarcity of land and water resources: A spatially explicit
641 mathematical programming approach. *Agric. Econ.* 39, 325–338. doi:10.1111/j.1574-
642 0862.2008.00336.x

643 Machovina, B., Feeley, K.J., 2014. Taking a bite out of biodiversity. *Science* (80-.). 343, 838 LP-
644 838.

645 Machovina, B., Feeley, K.J., Ripple, W.J., 2015. Biodiversity conservation: The key is reducing
646 meat consumption. *Sci. Total Environ.* 536, 419–431. doi:10.1016/j.scitotenv.2015.07.022

647 Marchal, V., Dellink, R., Van Vuuren, D., Clapp, C., Chateau, J., Magné, B., van Vliet, J., 2011. OECD
648 environmental outlook to 2050. *Organ. Econ. Co-operation Dev.*

649 Martín-Queller, E., Moreno-Mateos, D., Pedrocchi, C., Cervantes, J., Martínez, G., 2010. Impacts of
650 intensive agricultural irrigation and livestock farming on a semi-arid Mediterranean
651 catchment. *Environ. Monit. Assess.* 167, 423–435. doi:10.1007/s10661-009-1061-z

652 Mittermeier, R., Robles Gil, P., Hoffmann, M., Pilgrim, J., Brooks, T., Goettsch Mittermeier, C.,

653 Lamoreux, J., Fonseca, G., 2004. Hotspots revisited. earth's biologically richest and most
654 endangered terrestrial ecoregions., Conservation International.

655 Myers, N., Mittermeier, R.A., Mittermeier, C.G., Da Fonseca, G.A.B., Kent, J., 2000. Biodiversity
656 hotspots for conservation priorities. *Nature* 403, 853.

657 Naeem, S., Chazdon, R., Duffy, J.E., Prager, C., Worm, B., 2016. Biodiversity and human well-
658 being: an essential link for sustainable development. *Proc. R. Soc. B* 283, 20162091.

659 Naylor, R., Steinfeld, H., Falcon, W., Galloway, J., Smil, V., Bradford, E., Alder, J., Mooney, H., 2005.
660 Losing the links between livestock and land. *Science* (80-.). 310, 1621–1622.

661 Nelson, G.C., Valin, H., Sands, R.D., Havlík, P., Ahammad, H., Deryng, D., Elliott, J., Fujimori, S.,
662 Hasegawa, T., Heyhoe, E., Kyle, P., Von Lampe, M., Lotze-Campen, H., Mason d'Croz, D., van
663 Meijl, H., van der Mensbrugghe, D., Müller, C., Popp, A., Robertson, R., Robinson, S., Schmid,
664 E., Schmitz, C., Tabeau, A., Willenbockel, D., 2014. Climate change effects on agriculture:
665 economic responses to biophysical shocks. *Proc. Natl. Acad. Sci.* 111, 3274–9.
666 doi:10.1073/pnas.1222465110

667 Newbold, T., Hudson, L.N., Hill, S.L.L., Contu, S., Lysenko, I., Senior, R.A., Börger, L., Bennett, D.J.,
668 Choimes, A., Collen, B., Day, J., De Palma, A., Díaz, S., Echeverria-Londoño, S., Edgar, M.J.,
669 Feldman, A., Garon, M., Harrison, M.L.K., Alhusseini, T., Ingram, D.J., Itescu, Y., Kattge, J.,
670 Kemp, V., Kirkpatrick, L., Kleyer, M., Correia, D.L.P., Martin, C.D., Meiri, S., Novosolov, M.,
671 Pan, Y., Phillips, H.R.P., Purves, D.W., Robinson, A., Simpson, J., Tuck, S.L., Weiher, E., White,
672 H.J., Ewers, R.M., MacE, G.M., Scharlemann, J.P.W., Purvis, A., 2015. Global effects of land use
673 on local terrestrial biodiversity. *Nature* 520, 45–50. doi:10.1038/nature14324

674 Nijssen, M.E., WallisDeVries, M.F., Siepel, H., 2017. Pathways for the effects of increased nitrogen
675 deposition on fauna. *Biol. Conserv.* 212, 423–431.
676 doi:https://doi.org/10.1016/j.biocon.2017.02.022

677 O'Neill, B.C., Kriegler, E., Ebi, K.L., Kemp-Benedict, E., Riahi, K., Rothman, D.S., van Ruijven, B.J.,
678 van Vuuren, D.P., Birkmann, J., Kok, K., 2015. The roads ahead: narratives for shared
679 socioeconomic pathways describing world futures in the 21st century. *Glob. Environ.*
680 *Chang.*

681 O'Neill, B.C., Kriegler, E., Riahi, K., Ebi, K.L., Hallegatte, S., Carter, T.R., Mathur, R., van Vuuren,
682 D.P., 2014. A new scenario framework for climate change research: The concept of shared
683 socioeconomic pathways. *Clim. Change* 122, 387–400. doi:10.1007/s10584-013-0905-2

684 Orme, C.D.L., Davies, R.G., Burgess, M., Eigenbrod, F., Pickup, N., Olson, V.A., Webster, A.J., Ding,
685 T.-S., Rasmussen, P.C., Ridgely, R.S., 2005. Global hotspots of species richness are not
686 congruent with endemism or threat. *Nature* 436, 1016.

687 Pachauri, R.K., Allen, M.R., Barros, V.R., Broome, J., Cramer, W., Christ, R., Church, J.A., Clarke, L.,
688 Dahe, Q., Dasgupta, P., 2014. Climate change 2014: synthesis report. Contribution of
689 Working Groups I, II and III to the fifth assessment report of the Intergovernmental Panel
690 on Climate Change. IPCC.

691 Pereira, H.M., Navarro, L.M., Martins, I.S., 2012. Global biodiversity change: The bad, the good,
692 and the unknown. *Annu. Rev. Environ. Resour.* 37, 25–50. doi:10.1146/annurev-environ-
693 042911-093511

694 Pimm, S.L., Jenkins, C.N., Abell, R., Brooks, T.M., Gittleman, J.L., Joppa, L.N., Raven, P.H., Roberts,
695 C.M., Sexton, J.O., 2014. The biodiversity of species and their rates of extinction,
696 distribution, and protection. *Science* (80-.). 344. doi:10.1126/science.1246752

697 Pimm, S.L., Russell, G.J., Gittleman, J.L., Brooks, T.M., 1995. The future of biodiversity. *Science*
698 (80-.). 269, 347 LP-350.

699 Poore, J., Nemecek, T., 2018. Reducing food's environmental impacts through producers and
700 consumers. *Science* (80-.). 360, 987 LP-992.

701 Popp, A., Calvin, K., Fujimori, S., Havlik, P., Humpenöder, F., Stehfest, E., Bodirsky, B.L., Dietrich,
702 J.P., Doelmann, J.C., Gusti, M., Hasegawa, T., Kyle, P., Obersteiner, M., Tabeau, A., Takahashi,
703 K., Valin, H., Waldhoff, S., Weindl, I., Wise, M., Kriegler, E., Lotze-Campen, H., Fricko, O.,
704 Riahi, K., van Vuuren, D.P., 2016. Land-use futures in the shared socio-economic pathways.
705 *Glob. Environ. Chang.* 42, 331–345.
706 doi:<http://dx.doi.org/10.1016/j.gloenvcha.2016.10.002>

707 Popp, A., Lotze-Campen, H., Bodirsky, B., 2010. Food consumption, diet shifts and associated
708 non-CO2 greenhouse gases from agricultural production. *Glob. Environ. Chang.* 20, 451–
709 462. doi:10.1016/j.gloenvcha.2010.02.001

710 Possingham, H.P., Wilson, K.A., 2005. Biodiversity: Turning up the heat on hotspots. *Nature* 436,
711 919.

712 Pringle, R.M., 2017. Upgrading protected areas to conserve wild biodiversity. *Nature* 546, 91–99.
713 doi:10.1038/nature22902

714 Reich, P.B., 2009. Elevated CO2 reduces losses of plant diversity caused by nitrogen deposition.
715 *Science* (80-.). 326, 1399 LP-1402. doi:10.1126/science.1178820

716 Ren, G., Young, S.S., Wang, L., Wang, W., Long, Y., Wu, R., Li, J., Zhu, J., Yu, D.W., 2015.
717 Effectiveness of China's National Forest Protection Program and nature reserves. *Conserv.*
718 *Biol.* 29, 1368–1377. doi:10.1111/cobi.12561

719 Ripple, W.J., Estes, J.A., Beschta, R.L., Wilmers, C.C., Ritchie, E.G., Hebblewhite, M., Berger, J.,
720 Elmhagen, B., Letnic, M., Nelson, M.P., Schmitz, O.J., Smith, D.W., Wallach, A.D., Wirsing, A.J.,
721 2014a. Status and ecological effects of the world's largest carnivores. *Science* (80-.). 343.

722 Ripple, W.J., Smith, P., Haberl, H., Montzka, S.A., McAlpine, C., Boucher, D.H., 2014b. Ruminants,
723 climate change and climate policy. *Nat. Clim. Chang.* 4, 2–5. doi:10.1038/nclimate2081

724 Smith, B., Wärlind, D., Arneth, a., Hickler, T., Leadley, P., Siltberg, J., Zaehle, S., 2014. Implications
725 of incorporating N cycling and N limitations on primary production in an individual-based
726 dynamic vegetation model. *Biogeosciences* 11, 2027–2054. doi:10.5194/bg-11-2027-2014

727 Stehfest, E., Bouwman, L., Van Vuuren, D.P., Den Elzen, M.G.J., Eickhout, B., Kabat, P., 2009.
728 Climate benefits of changing diet. *Clim. Change* 95, 83–102.

729 Stevens, C.J., Dise, N.B., Mountford, J.O., Gowing, D.J., 2004. Impact of Nitrogen Deposition on the
730 Species Richness of Grasslands. *Science* (80-.). 303, 1876 LP-1879.
731 doi:10.1126/science.1094678

732 Stoll-Kleemann, S., Schmidt, U.J., 2017. Reducing meat consumption in developed and transition
733 countries to counter climate change and biodiversity loss: a review of influence factors.
734 *Reg. Environ. Chang.* 17, 1261–1277. doi:10.1007/s10113-016-1057-5

735 Swain, M., Blomqvist, L., McNamara, J., Ripple, W.J., 2018. Reducing the environmental impact of
736 global diets. *Sci. Total Environ.* 610, 1207–1209.

737 Taylor, K.E., Stouffer, R.J., Meehl, G.A., 2012. An overview of CMIP5 and the experiment design.
738 *Bull. Am. Meteorol. Soc.* 93, 485–498.

739 Ten Brink, B., van der Esch, S., Kram, T., Van Oorschot, M., Arets, E., 2010. Rethinking Global
740 Biodiversity Strategies: Exploring structural changes in production and consumption to
741 reduce biodiversity loss. Netherlands Environmental Assessment Agency.

742 Tilman, D., Balzer, C., Hill, J., Befort, B.L., 2011. Global food demand and the sustainable

intensification of agriculture. *Proc. Natl. Acad. Sci.* 108, 20260–4.
doi:10.1073/pnas.1116437108

Tilman, D., Clark, M., 2014a. Global diets link environmental sustainability and human health. *Nature* 515, 518–522.

Tilman, D., Clark, M., 2014b. Global diets link environmental sustainability and human health. *Nature* 515, 518–522.
doi:http://www.nature.com/nature/journal/v515/n7528/full/nature13959.html

Tilman, D., Clark, M., Williams, D.R., Kimmel, K., Polasky, S., Packer, C., 2017. Future threats to biodiversity and pathways to their prevention. *Nature* 546, 73–81.
doi:10.1038/nature22900

Tilman, D., Fargione, J., Wolff, B., D'Antonio, C., Dobson, A., Howarth, R., Schindler, D., Schlesinger, W.H., Simberloff, D., Swackhamer, D., 2001. Forecasting agriculturally driven global environmental change. *Science* (80-.). 292, 281 LP-284.

UNEP-WCMC., I&, 2016. The World Database on Protected Areas (WDPA). [WWW Document].

van Vuuren, D.P., Edmonds, J., Kainuma, M., Riahi, K., Thomson, A., Hibbard, K., Hurtt, G.C., Kram, T., Krey, V., Lamarque, J.F., Masui, T., Meinshausen, M., Nakicenovic, N., Smith, S.J., Rose, S.K., 2011. The representative concentration pathways: An overview. *Clim. Change* 109, 5–31.
doi:10.1007/s10584-011-0148-z

Visconti, P., Bakkenes, M., Baisero, D., Brooks, T., Butchart, S.H.M., Joppa, L., Alkemade, R., Di Marco, M., Santini, L., Hoffmann, M., Maiorano, L., Pressey, R.L., Arponen, A., Boitani, L., Reside, A.E., van Vuuren, D.P., Rondinini, C., 2016. Projecting global biodiversity indicators under future development scenarios. *Conserv. Lett.* 9, 5–13. doi:10.1111/conl.12159

Visconti, P., Bakkenes, M., Smith, R.J., Joppa, L., Sykes, R.E., 2015. Socio-economic and ecological impacts of global protected area expansion plans. *Philos. Trans. R. Soc. B Biol. Sci.* 370, 20140284. doi:10.1098/rstb.2014.0284

Wellesley, L., Happer, C., Froggatt, A., 2015. Changing climate, changing diets: pathways to lower meat consumption. Royal Institute of International Affairs, Chatham House.

Westhoek, H., Rood, T., van den Berg, M., Janse, J., Nijdam, D., Reudink, M., Stehfest, E., Lesschen, J.P., Oenema, O., Woltjer, G.B., 2011. The protein puzzle: the consumption and production of meat, dairy and fish in the European Union. Netherlands Environmental Assessment Agency.

Wirsén, S., Azar, C., Berndes, G., 2010. How much land is needed for global food production under scenarios of dietary changes and livestock productivity increases in 2030? *Agric. Syst.* 103, 621–638.

Yamaguchi, T., Blumwald, E., 2005. Developing salt-tolerant crop plants: challenges and opportunities. *Trends Plant Sci.* 10, 615–620.